

Photoelectrochemical Hydrogen Production (Project ID: PD057)

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2010 DOE Hydrogen Program

Washington DC, June 7, 2010

Project ID PD057

Overview

Timeline

- July 2006
- August 2010
- Percent complete 80%

Budget

- Total project funding \$
 - **Hydrogen Production \$297K**
 - **Contractor' share \$127,181**
 - DOE Total share \$ 890,998DOE share for Hydrogen Storage \$594K; Contractor \$254,362.
- Total funding received in FY08 \$ 324,721(for generation and storage)

Barriers

- Barriers addressed
 - (AP) Materials Efficiency
 - (AQ) Materials Durability
 - (AS) Device Configuration Designs

Partners

- Interactions/ collaborations
 1. University of Nevada, Reno
 2. Arkansas Nanotechnology Center, University of Arkansas at Little Rock
 3. NASA Kennedy Space Center
 4. Boston University

Objectives

Overall	Optimize surface properties of photoanodes for efficient generation of Hydrogen
2007-08	<ul style="list-style-type: none">• Study interfacial charge transfer process in PEC• Investigate the roles of surface states and nano-structures of Photoanodes (Use TiO₂ electrode as a model)• Develop partnership with other institutions• Develop outreach and educational programs
2008-09	<ul style="list-style-type: none">• Remove surface contaminants & charge traps by He plasma treatments• Surface doping of TiO₂ electrode with N using plasma treatment; Create oxygen vacancies• Improve light absorption cross section and interfacial charge transfer rate with nanostructured anodes• Measure photocurrent density of nano-structured TiO₂ electrodes with simulated solar radiation

Objectives

Overall	Optimize nanostructures of TiO₂ Photoanodes for light absorption and corrosion resistance
2009-2010	<ul style="list-style-type: none">• Optimize the structure of the nanotubes by varying anodization voltage• Use layered electrodes of TiSi₂ and TiO₂,• Optimize interfacial charge transfer,• Develop processes for synthesizing layered patterned nanotubular electrodes of TiSi₂ and TiO₂

Rationale

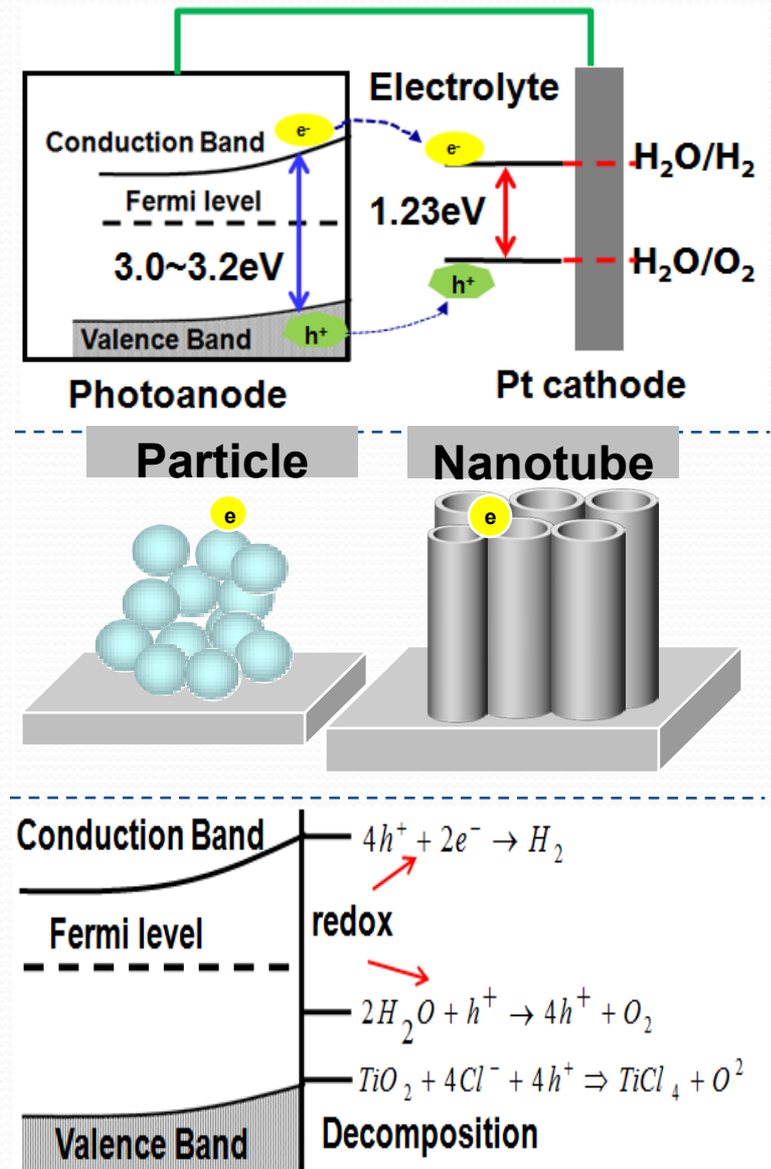
- Optimize interfacial processes: photon absorption, charge separation, minimization of carrier traps.
- No single semiconductor electrode has been found to provide both high absorption and corrosion resistance
- TiSi_2 (bandgap ranging from 3.4 eV to 1.5 eV) can harvest the entire visible and UV spectrum of solar radiation. TiO_2 electrode layer can be used to minimize photocorrosion.

Approach

- Remove surface contaminants and surface states that act as charge carrier traps
- Apply Plasma surface modification for surface cleaning as well as for surface doping of n-type dopants (N)
- Test surface modified nanostructured TiSi_2 and TiO_2 anodes for photoelectrochemical generation of hydrogen

TiO₂ photoanode nanotube

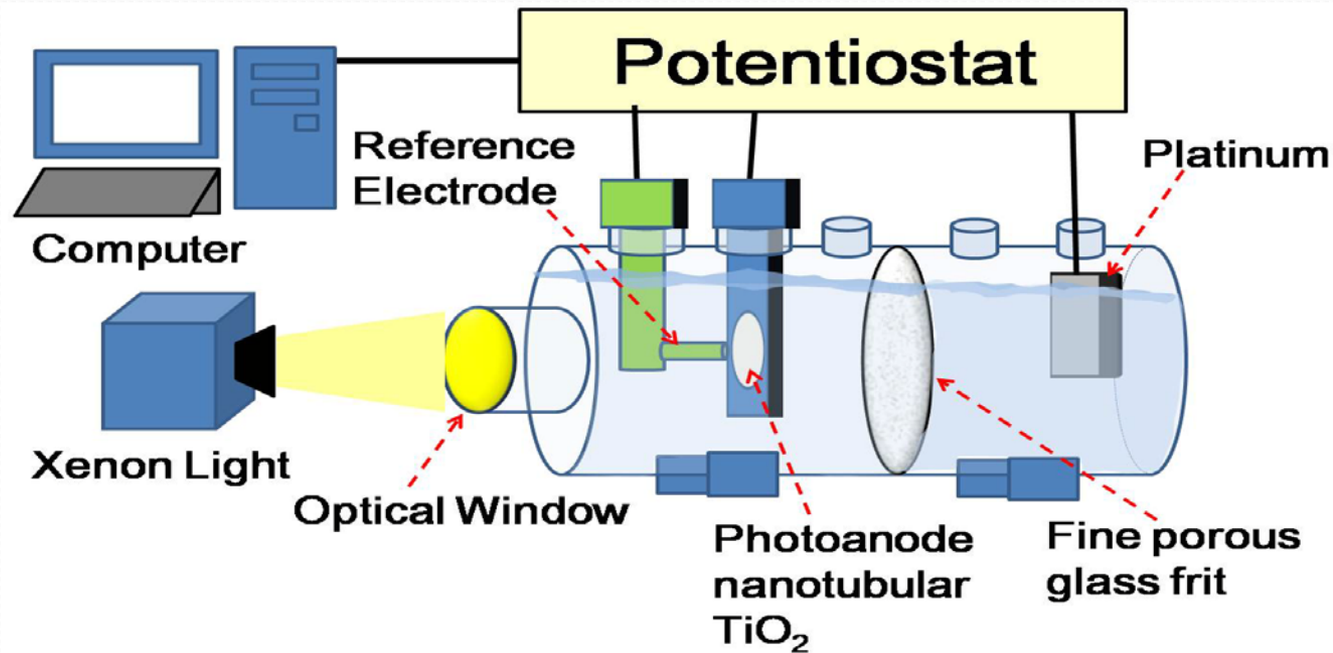
- Energy band gap required more than 1.23eV
 - TiO₂ energy band gap is 3.0~3.2eV (7% of the solar spectrum)
- TiO₂ nanotube array (electron mobility is higher compared to that of nanoparticle photoanodes)
- Durability (decomposition is slow)
- Non toxic



Experimental procedure

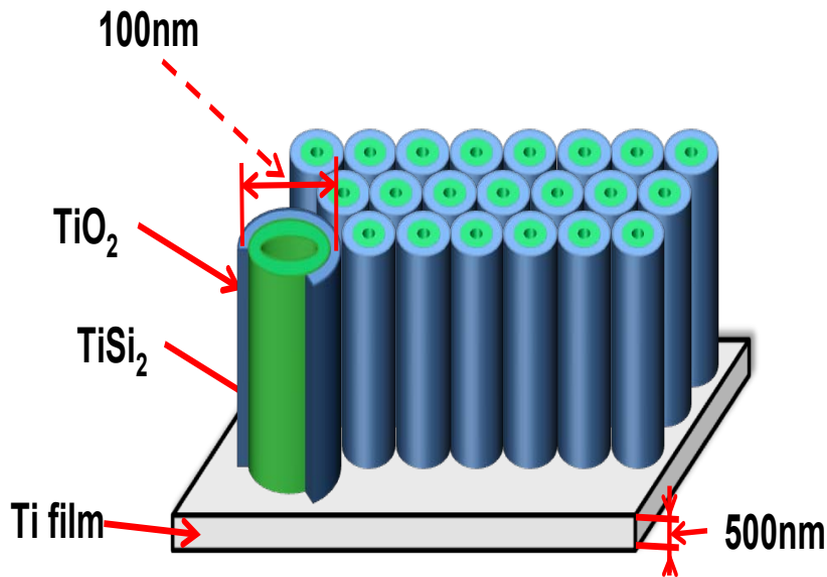
- Electrochemical synthesis of TiO_2 nanotube array by anodization of Ti film
- Plasma surface modification of TiO_2 nanorods using Nitrogen plasma
- Surface structure measurements
- Photocurrent density measurements

Photoelectrochemical test of photoanode



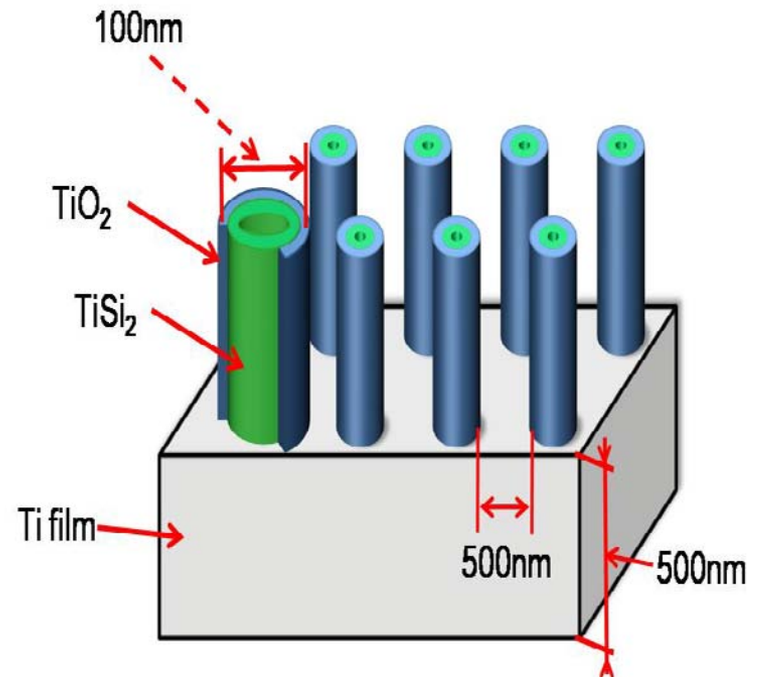
- Potentiostat/Galvanostat model 283.
- Xenon lamp (30 mW)
- 60 mm diameter quartz optical window
- A reference electrode (Ag/AgCl) was placed close to the anode
- Electrolyte; 1M KOH (pH~14) + DI water solution

Nanotubular Layered Photoelectrode



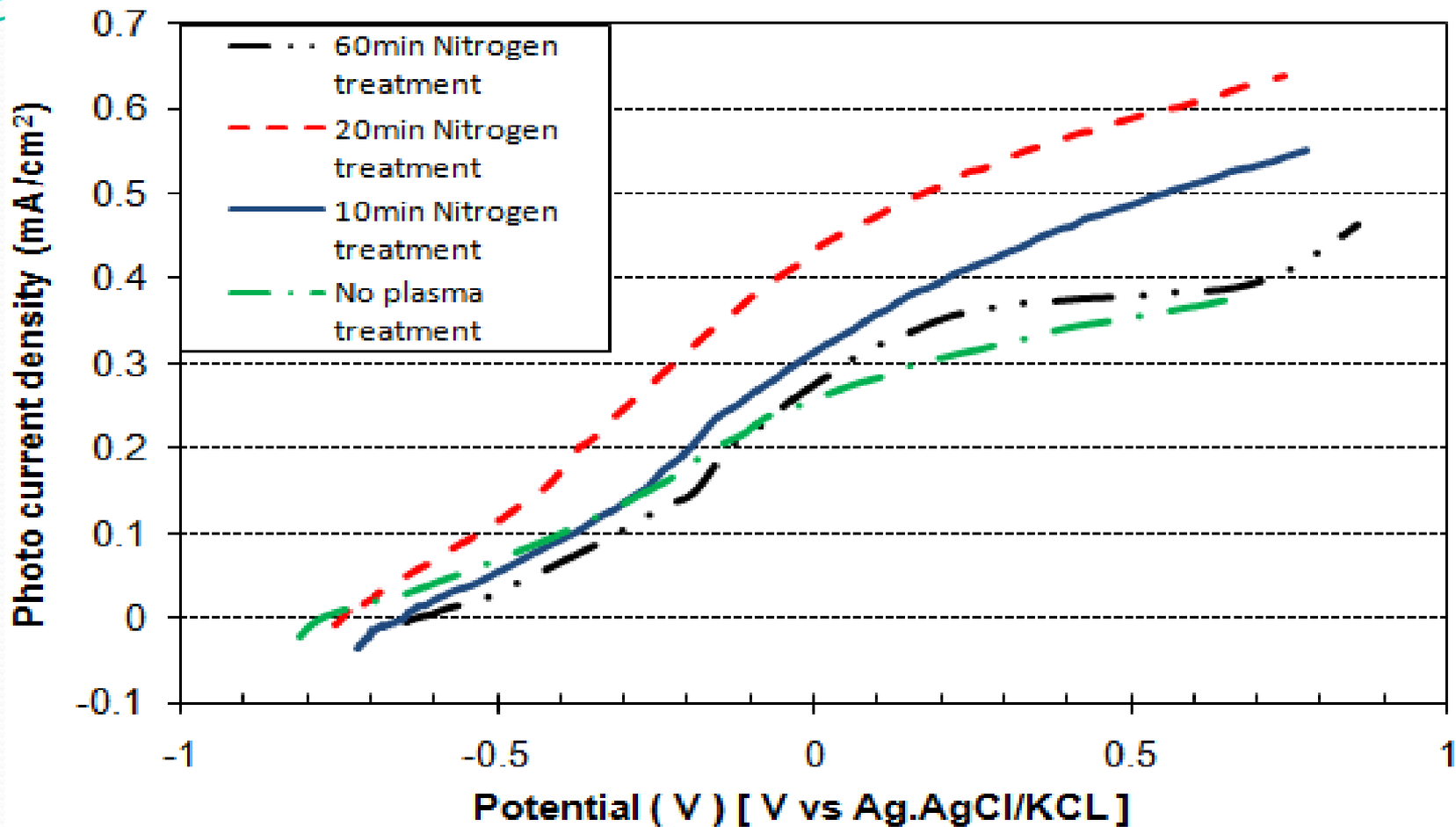
Close Spacing

Patterned nanotubular structure for maximizing light absorption



Open spacing

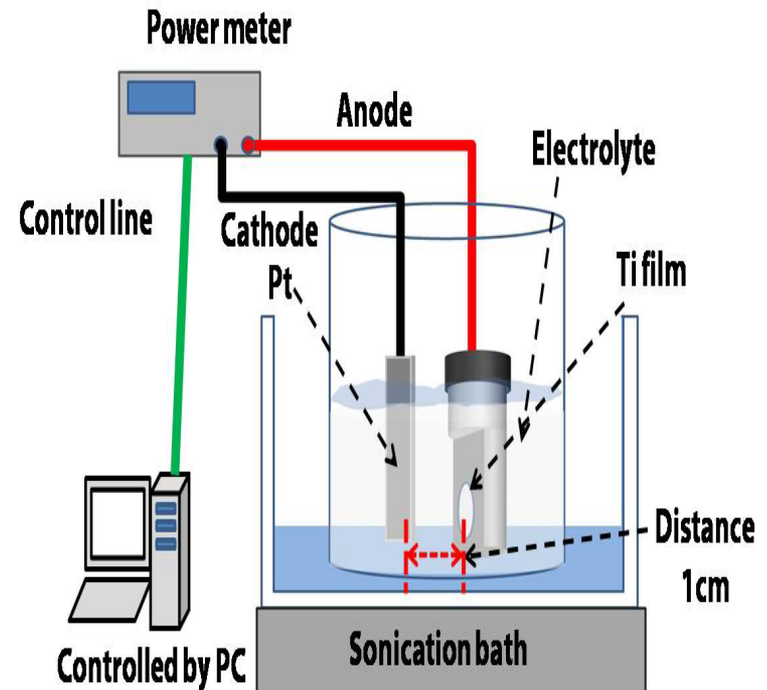
Result- Photocurrent vs. bias voltage



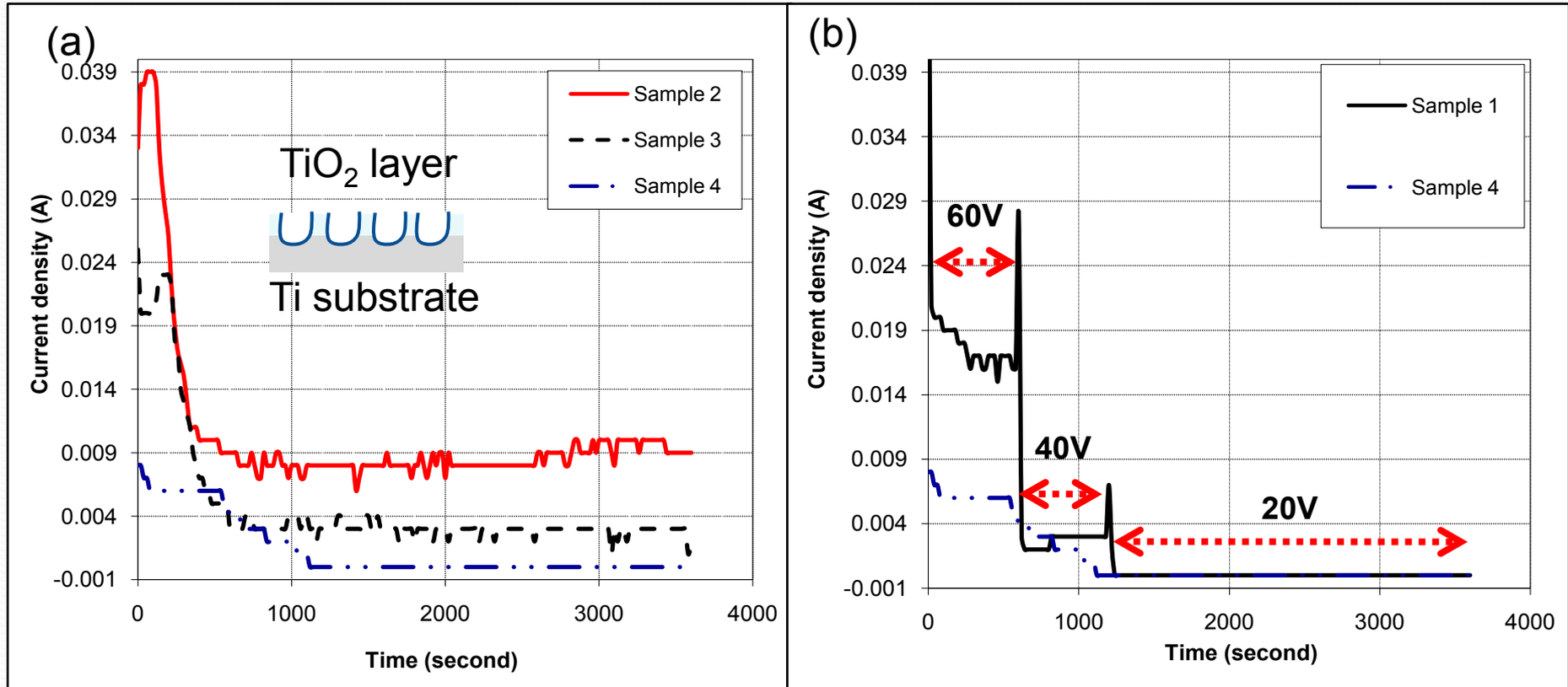
- Photocurrent density vs. bias voltage plotted for anodized samples after plasma treatment for different time periods in nitrogen atmosphere. Anodization was performed at 20V for 60 min and all samples were annealed

Equipment for anodization

- Anodization
 - The dimensions of the nanotubes can be easily varied (Length, wall thickness and internal diameter)
- Two-electrode electrochemical cell
- The electrolyte consisted of 0.5 wt % NH_4F + ethylene glycol + 0.2 % by vol DI water
- Room temperature anodization at a constant potential

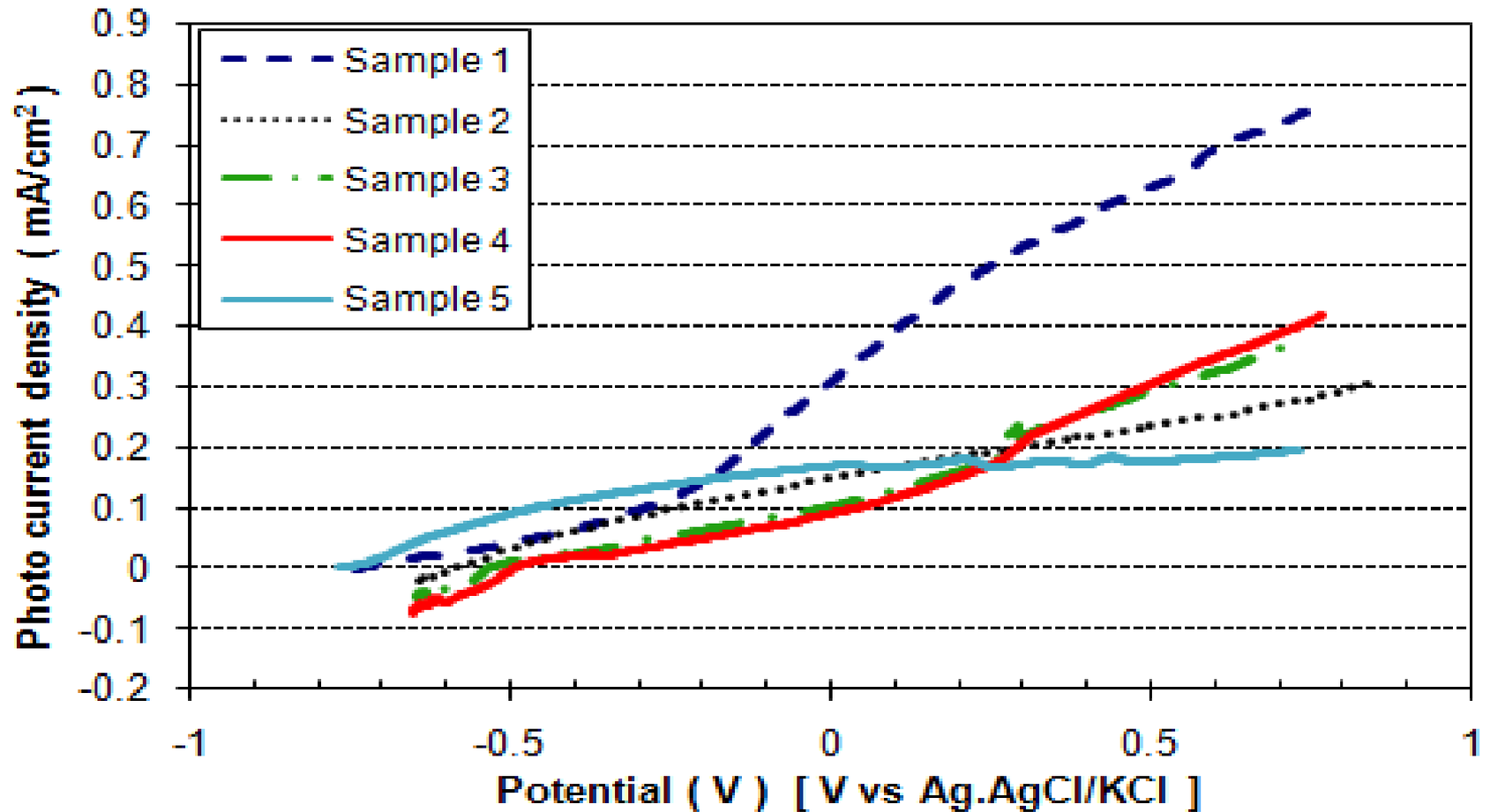


Results-current transients during anodization



- Current density vs time during anodization; (a) constant voltage anodization at different voltage levels, (b) step voltage anodization at 60V-40V-20V compared with constant 20 V anodization

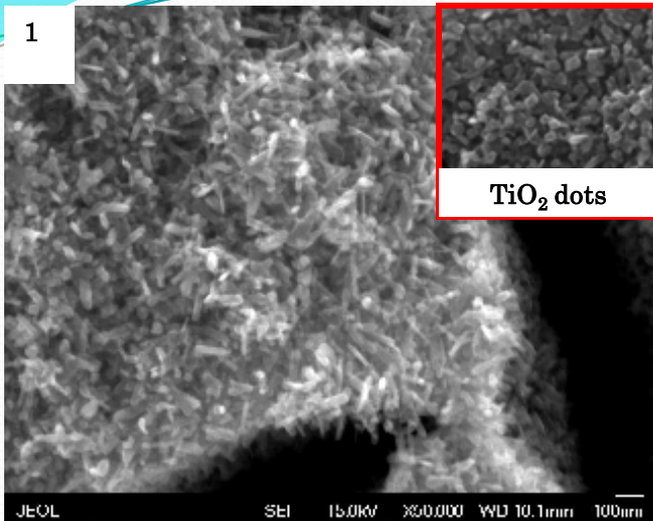
Photocurrent density vs. anode nanostructure



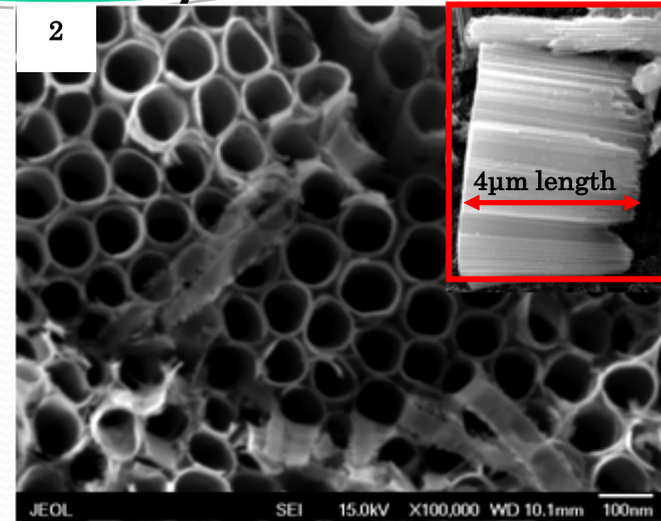
Photocurrent density vs bias voltage plotted for samples anodized at different voltages

Stepped voltage anodization resulted in 55% higher current compared to the value of photocurrent at a constant voltage anodization

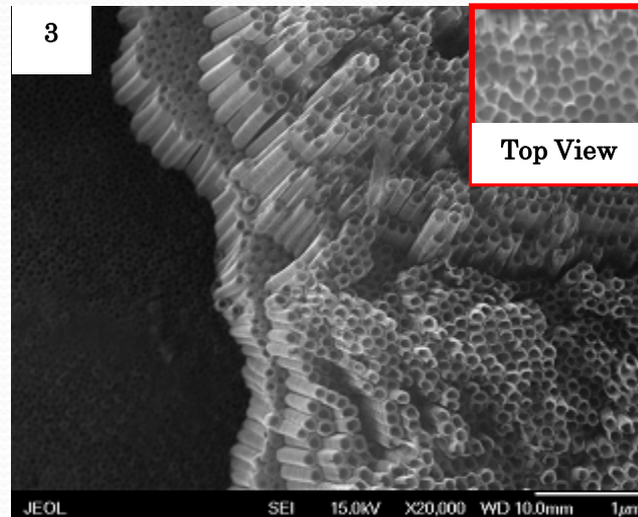
SEM image of TiO₂ nanotubular array



- The nanotube;
 - Average length 4 μm
 - Average diameter 100 nm
 - Average wall thickness 5 nm
 - Average center to center spacing of 150 nm



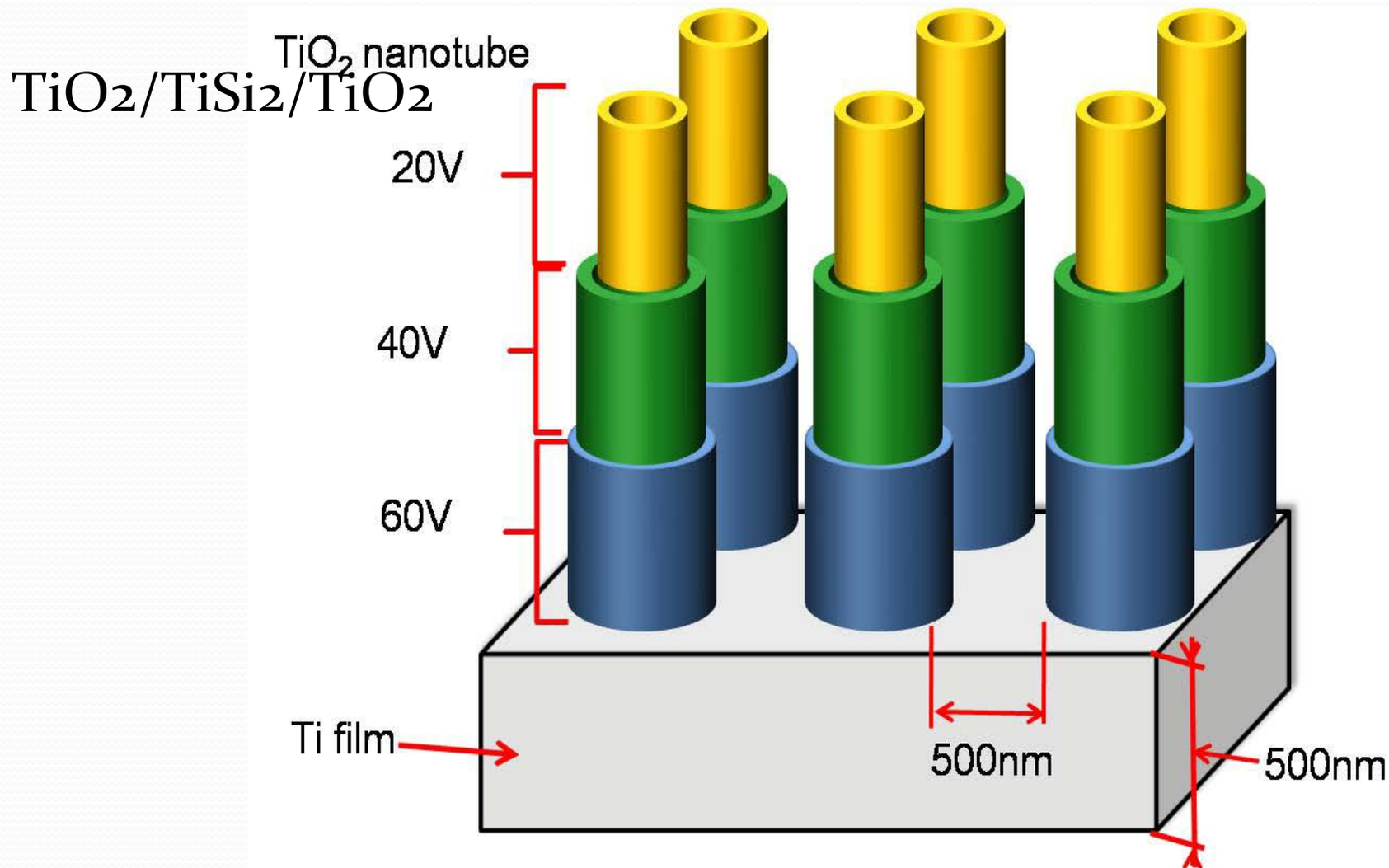
- The nanorod;
 - Average length 100nm
 - Average width 10nm
 - Average height 10nm
- The nanodot
 - Average 10 nm of square of side.



- The three successive stage nanotube;
 - Average length 4 μm
 - Average diameter 100nm
 - Average wall thickness 5nm
 - Average center to center spacing of 150 nm

SEM image showing the TiO₂ sample anodized in a fluoride based electrolyte at three different voltage settings: (1) constant potential 20V (Fig.2a), (2) constant potential 60V (Fig.2b), and (3) stepped voltage potentials, 60V-40V-20V (Fig.2c)

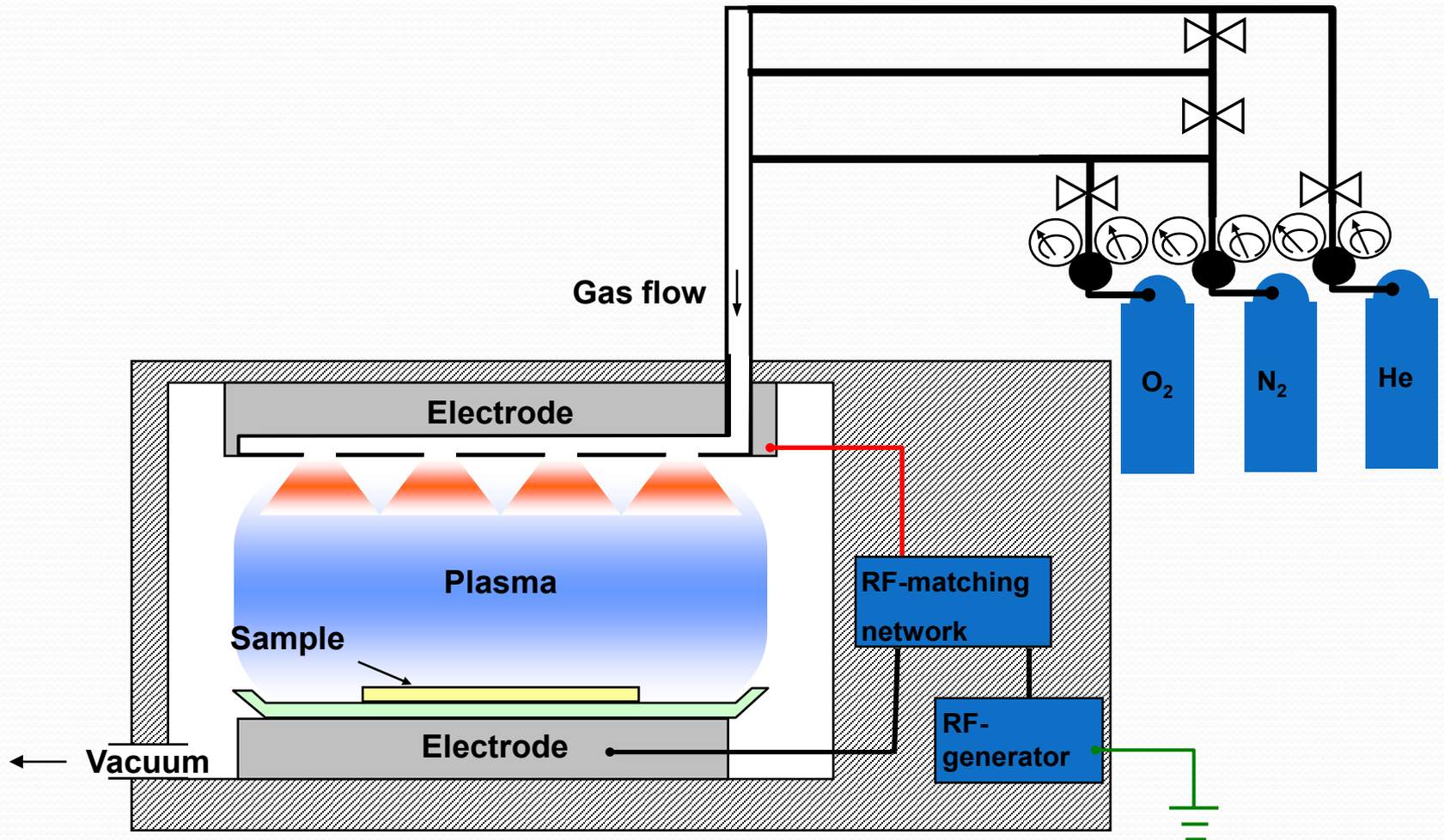
Layered Nanotubular Arrays



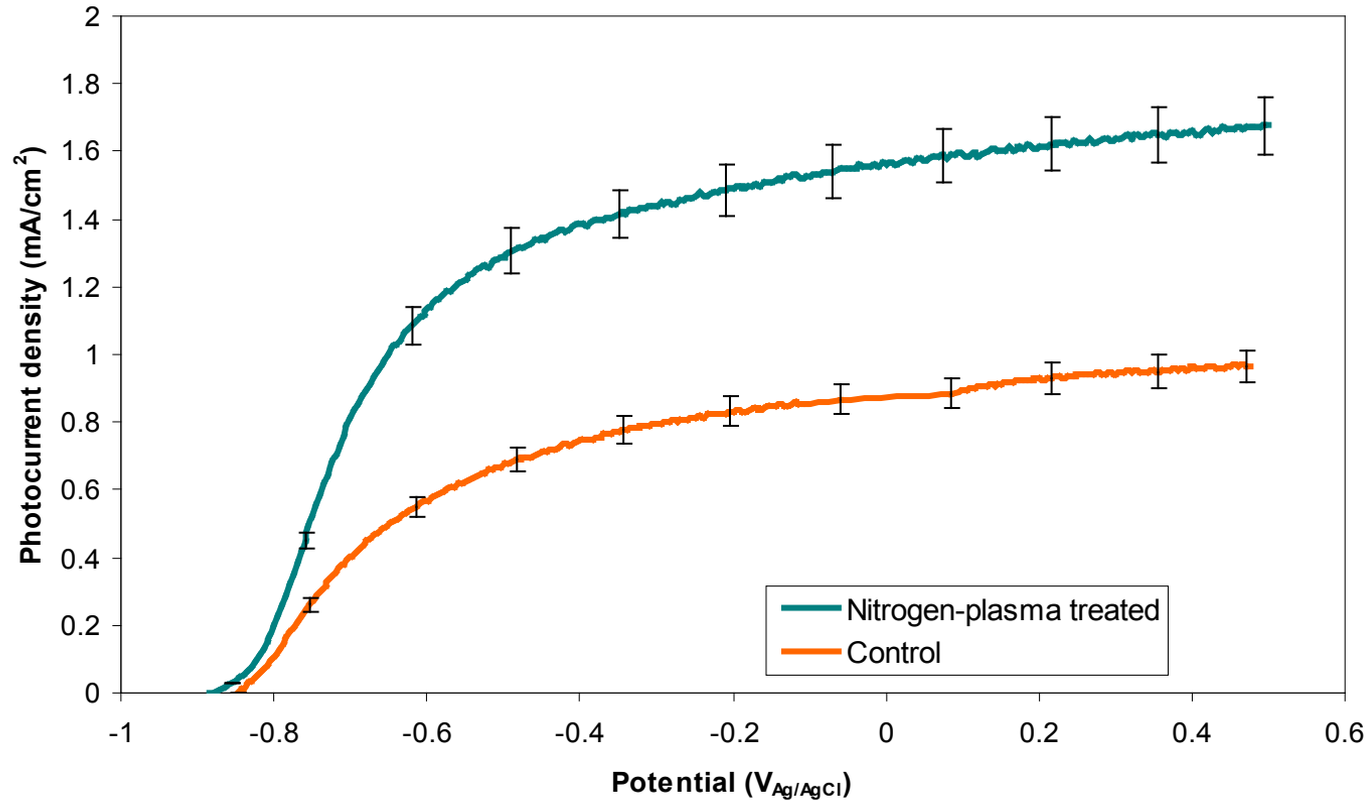
Plasma surface modification

- Plasma surface modification was performed using low-pressure Nitrogen plasma [13.56 MHz rf 200 W plasma at an operating pressure 150 mtorr]
- Samples were exposed to plasma for 10 minutes in each test run
- Untreated and nitrogen plasma treated samples were tested for photocurrent density

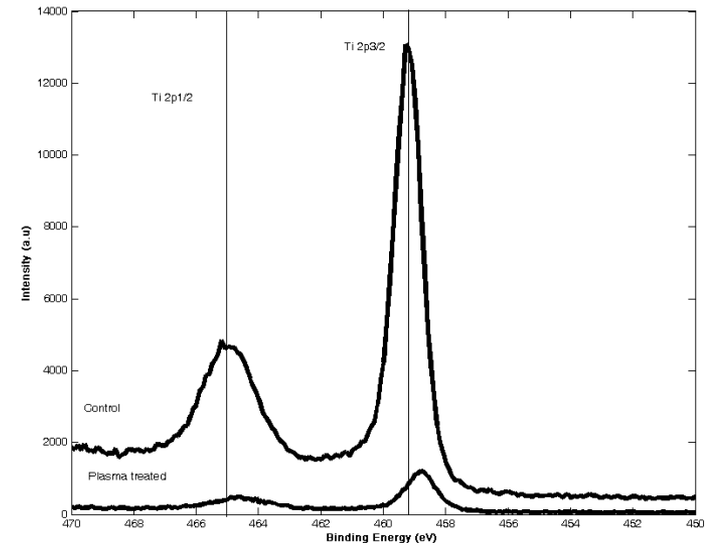
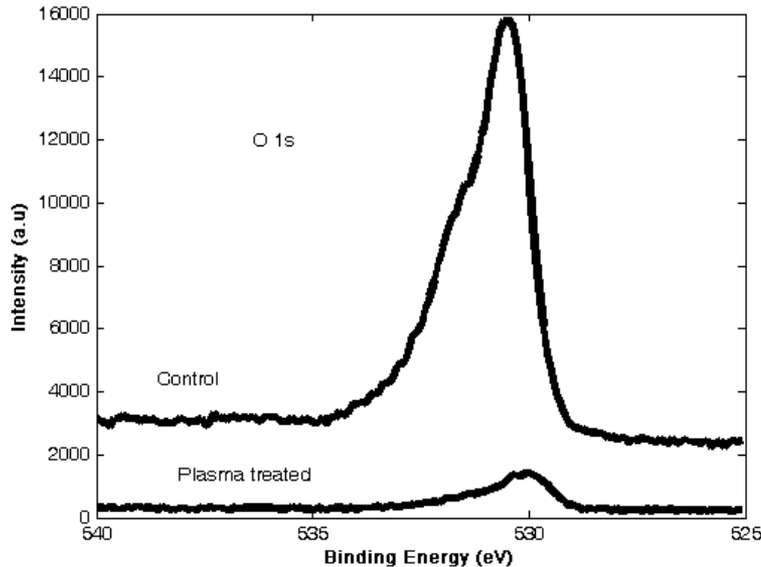
Schematic of low-pressure plasma reactor used for surface modification



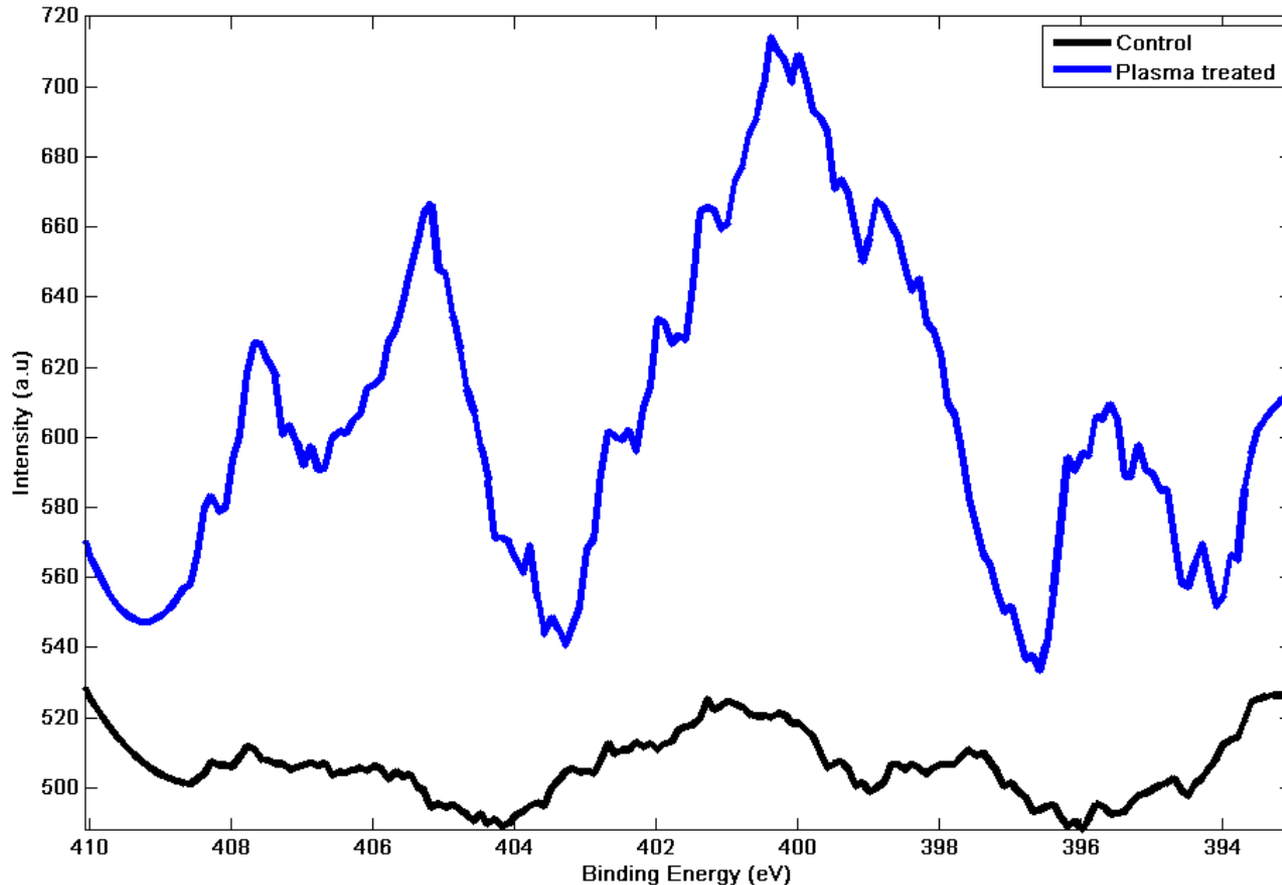
Photocurrent density measurements for control and nitrogen plasma treated titania photoanodes



XPS Analysis of Nitrogen plasma treated TiO₂ samples



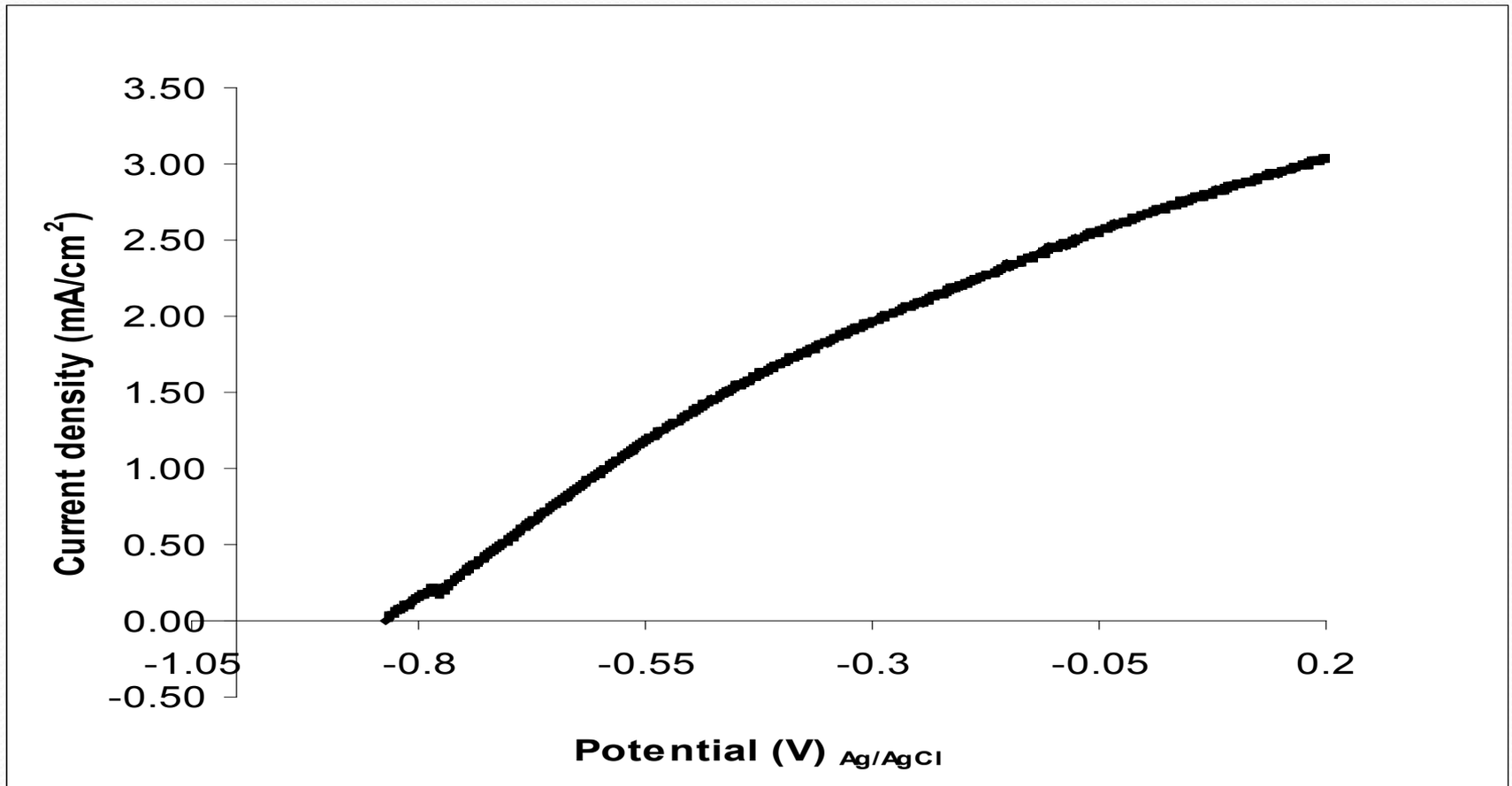
XPS spectrum of (a) Ti 2p and (b) O 1s levels for Control and Nitrogen-Plasma treated TiO₂ photoanodes



XPS spectrum (a) Control and (b) Nitrogen-Plasma treated TiO₂ photoanodes

The narrow scan N 1s spectrum is demonstrated in peaks at 400 and 396 eV, which have been ascribed to presence of nitrogen in lattice structure either as substitutional dopant for O, or as interstitial dopant

Photocurrent Density with TiSi₂ Particles on TiO₂ Nanotubular Electrode



Effects of Surface and Structural Modifications of TiO₂ Nanotubes

Table 1. Enhancements of open circuit potential and photocurrent density values by surface modification of oxygen-annealed samples by nitrogen plasma treatment.

Sample description	OCP (bright), $V_{Ag/AgCl}$	Current at -0.2 V (mA cm^{-2})	Current at 0.2 V (mA cm^{-2})
Control	-0.85	0.83	0.93
N ₂ plasma treated	-0.97	0.94	1.68

Table 2. Structural Modifications of nanotubular TiO₂ photoanodes by changing anodization conditions of Ti foil (Stepped voltage (Sample 1) vs. constant voltage anodization)

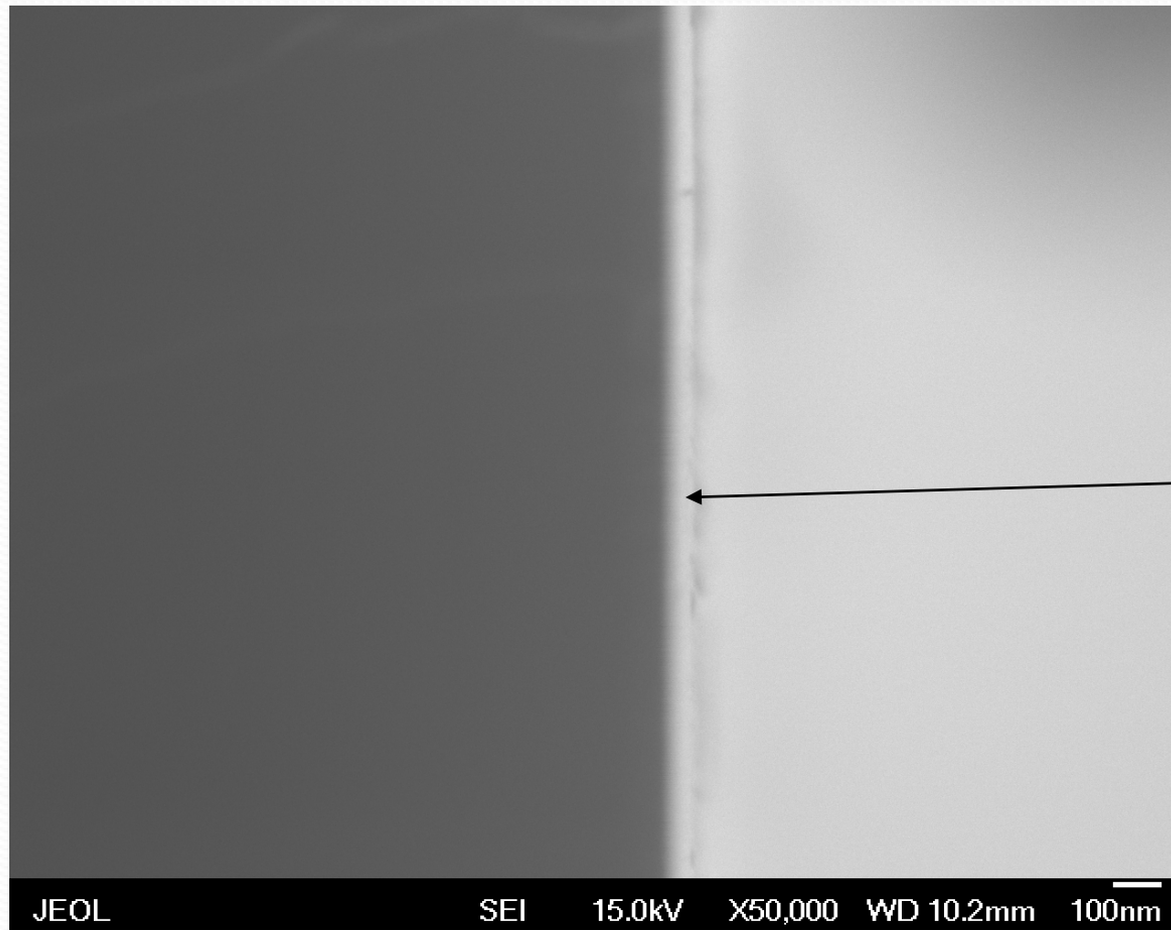
Sample Name	Anodization Voltage and duration		
Sample 1	60 V for 10 min	40 V for 10 min	20 V for 40min
Sample 2	60 V for 60 min		
Sample 3	40 V for 60 min		
Sample 4	20 V for 60 min		
Sample 5	20 V for 60 min		

Synthesis of Titanium disilicide thin films for using composite multilayer photoanodes for Hydrogen generation

- Deposit a thin film of TiSi_2 using sputter coating on Ti film for 60 minutes
- Analyze chemical composition gradients using XPS and EDS
- Use electron beams, impinged with different energies (3, 7, 10, 15, and 30 kV) on the film to analyze chemical composition at different depths of the film

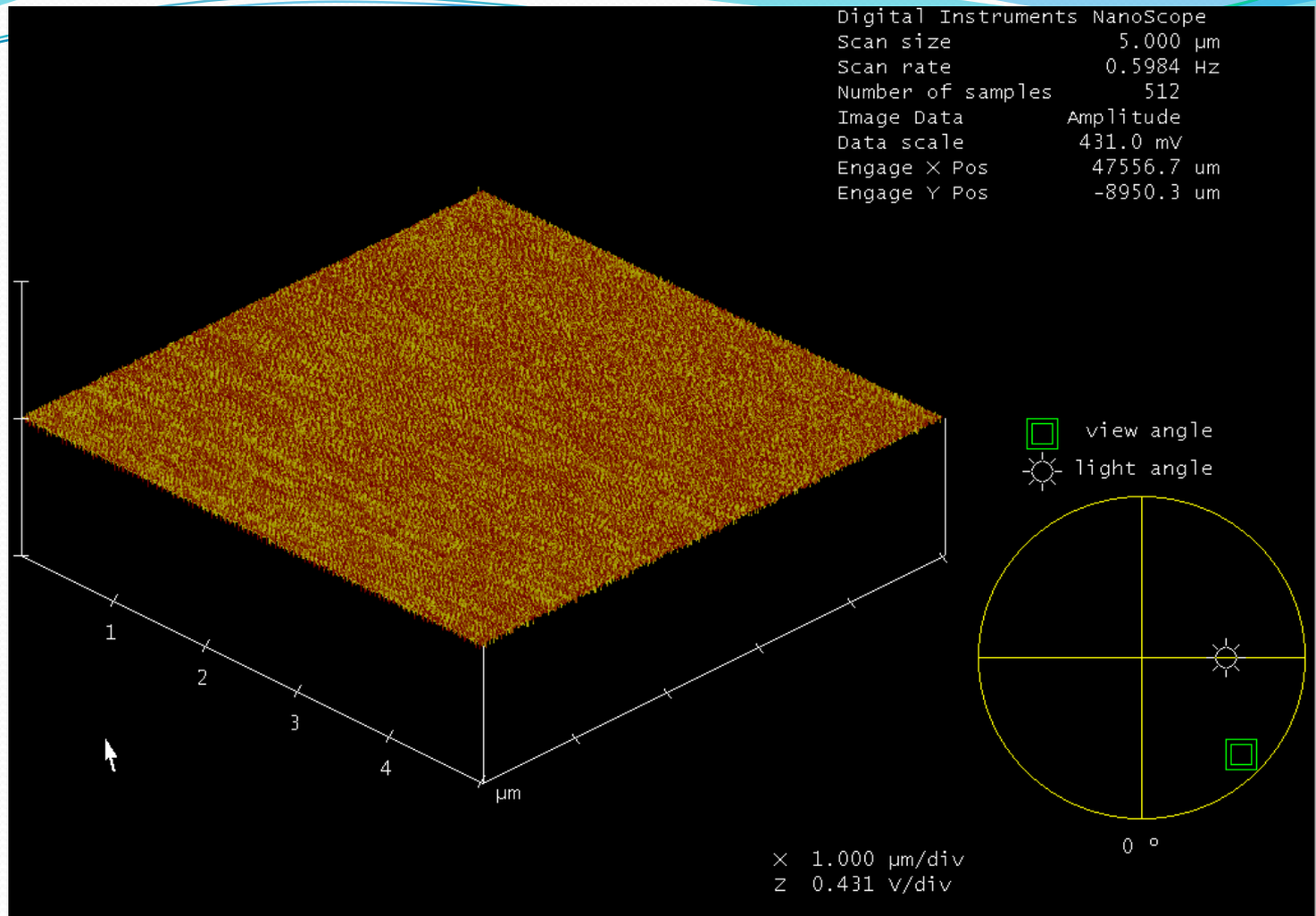
Development of Heterojunction $\text{TiSi}_2/\text{TiO}_2$ photoanodes

- Titanium disilicide is a promising photoanode material in photoelectrochemical hydrogen generation
- A heterojunction $\text{TiO}_2/\text{TiSi}_2$ photoanode can harvest a significant portion of solar radiation in the visible region.
- Broadband reflectance measurements for TiSi_2 show a bandgap ranging from 3.4 eV to 1.5 eV. However TiSi_2 is unstable in water.
- TiO_2 film coatings can serve as a protective layer for TiSi_2 .



← Titanium silicide film

SEM image of a titanium silicide film



AFM image of a titanium silicide film displaying high degree of uniformity

Results

- **Nitrogen plasma treatment of titania photoanodes resulted in 80% increase in photocurrent density;**
- **XPS analysis clearly indicated the incorporation of N in titania lattice structure.**
- **Electrochemical synthesis of titania nanotube arrays using needle shaped cathode improved the performance of photoanode**
- **The deposition rate of titanium silicide has been determined as a part of the process of developing heterojunction $\text{TiO}_2/\text{TiSi}_2$ photoanodes**

Technical Accomplishments

- **Plasma surface modification and surface doping increased photocurrent density of titania nanotubular electrodes**
- **Synthesis of TiO_2 nanorods**
- **Synthesis of TiSi_2 based heterojunction electrodes**
- **Photoelectrochemical hydrogen production
with a high photocurrent density**

Work under progress:

- **Develop patterned nanotubular layered TiSi_2 and TiO_2 photoanodes.**
- **Characterize interfacial surface states between TiSi_2 and TiO_2 photoanodes by determining density of states, surface bandgaps, crystallinity, optical absorption spectrum, photoconductivity, and durability.**
- **Optimize interface states and the process of layered photoanode synthesis**

Future Work

- **Develop multi-junction (TiSi₂ and TiO₂) electrodes to enhance the absorption of solar radiation**
- **Analyze the density of the interface states and the distribution of surface bandgaps**
- **Measure light absorption vs λ**
- **Measure photocurrent conversion efficiency (IPCE vs λ) and measure corrosion resistance**
- **Determine photo-generated carrier concentration decay by using a rf-conductivity probe**
- **Perform multi-dimensional analysis: cost, durability, efficiency and environmental impact in PEC based hydrogen generation**

Project Summary

Relevance: Develop efficient photoanode materials for optimizing hydrogen production

Approach: Plasma surface modification for removing surface contaminants and use of layered electrodes for PEC based generation of hydrogen

Technical Accomplishment and Progress: Enhanced photocurrent density with oxygen annealed photoanodes of TiSi_2 and TiO_2 photoanodes with surface doping of nitrogen using plasma treatments and synthesis of nanostructured electrodes

Collaboration: University of Nevada, Reno and Arkansas Nanotechnology Center

Proposed future research: Application of layered photoanodes, plasma treatments and quantitative determination of photoelectrochemical generation of hydrogen

Acknowledgements: Department of Energy, University of Nevada, Reno, NASA Kennedy Space Center, Arkansas Science and Technology Authority, and Boston University

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Project ID #PD057